

Connecting via Winsock to STN

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LOGINID:SSSPTA1642BJF

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TERMINAL (ENTER 1, 2, 3, OR ?):2

\*\*\*\*\* Welcome to STN International \*\*\*\*\*

NEWS	1		Web Page for STN Seminar Schedule - N. America
NEWS	2	JAN 02	STN pricing information for 2008 now available
NEWS	3	JAN 16	CAS patent coverage enhanced to include exemplified prophetic substances
NEWS	4	JAN 28	USPATFULL, USPAT2, and USPATOLD enhanced with new custom IPC display formats
NEWS	5	JAN 28	MARPAT searching enhanced
NEWS	6	JAN 28	USGENE now provides USPTO sequence data within 3 days of publication
NEWS	7	JAN 28	TOXCENTER enhanced with reloaded MEDLINE segment
NEWS	8	JAN 28	MEDLINE and LMEEDLINE reloaded with enhancements
NEWS	9	FEB 08	STN Express, Version 8.3, now available
NEWS	10	FEB 20	PCI now available as a replacement to DPCI
NEWS	11	FEB 25	IFIREF reloaded with enhancements
NEWS	12	FEB 25	IMSPRODUCT reloaded with enhancements
NEWS	13	FEB 29	WPINDEX/WPIDS/WPIX enhanced with ECLA and current U.S. National Patent Classification
NEWS	14	MAR 31	IFICDB, IFIPAT, and IFIUDB enhanced with new custom IPC display formats
NEWS	15	MAR 31	CAS REGISTRY enhanced with additional experimental spectra
NEWS	16	MAR 31	CA/Caplus and CASREACT patent number format for U.S. applications updated
NEWS	17	MAR 31	LPCI now available as a replacement to LDPCI
NEWS	18	MAR 31	EMBASE, EMBAL, and LEMBASE reloaded with enhancements
NEWS	19	APR 04	STN AnaVist, Version 1, to be discontinued
NEWS	20	APR 15	WPIDS, WPINDEX, and WPIX enhanced with new predefined hit display formats
NEWS	21	APR 28	EMBASE Controlled Term thesaurus enhanced
NEWS	22	APR 28	IMSRESEARCH reloaded with enhancements
NEWS	23	MAY 30	INPAFAMDB now available on STN for patent family searching
NEWS	24	MAY 30	DGENE, PCTGEN, and USGENE enhanced with new homology sequence search option
NEWS	25	JUN 06	EPFULL enhanced with 260,000 English abstracts
NEWS	26	JUN 06	KOREAPAT updated with 41,000 documents
NEWS	27	JUN 13	USPATFULL and USPAT2 updated with 11-character patent numbers for U.S. applications
NEWS	28	JUN 19	CAS REGISTRY includes selected substances from web-based collections
NEWS	29	JUN 25	CA/Caplus and USPAT databases updated with IPC reclassification data
NEWS	30	JUN 30	AEROSPACE enhanced with more than 1 million U.S. patent records
NEWS	31	JUN 30	EMBASE, EMBAL, and LEMBASE updated with additional options to display authors and affiliated



E7	1	EPOTHILONE B HYDROXYLASE/CN
E8	1	EPOTHILONE B HYDROXYLASE (AMYCOLATOPSIS ORIENTALIS GENE EBH)/CN
E9	1	EPOTHILONE B N-OXIDE/CN
E10	1	EPOTHILONE B10/CN
E11	1	EPOTHILONE C/CN
E12	1	EPOTHILONE C BIS(TERT-BUTYLDIMETHYLSILYL) ETHER/CN
E13	1	EPOTHILONE C/D 12,13-EPOXIDASE/CN
E14	1	EPOTHILONE C/D MONOOXYGENASE/CN
E15	1	EPOTHILONE C/D SYNTHETASE/CN
E16	1	EPOTHILONE C1/CN
E17	1	EPOTHILONE C2/CN
E18	1	EPOTHILONE C3/CN
E19	1	EPOTHILONE C4/CN
E20	1	EPOTHILONE C5/CN
E21	1	EPOTHILONE C6/CN
E22	1	EPOTHILONE C7/CN
E23	1	EPOTHILONE C8/CN
E24	1	EPOTHILONE C9/CN
E25	1	EPOTHILONE D/CN

=> S E3

L1 1 "EPOTHILONE B"/CN

=> S L1 EXA SAM

SAMPLE IS IGNORED AS A SCOPE FOR THIS SEARCH

L2 1 "EPOTHILONE B"/CN

=> DIS L2 1 SAM

THE ESTIMATED COST FOR THIS REQUEST IS 1.04 U.S. DOLLARS

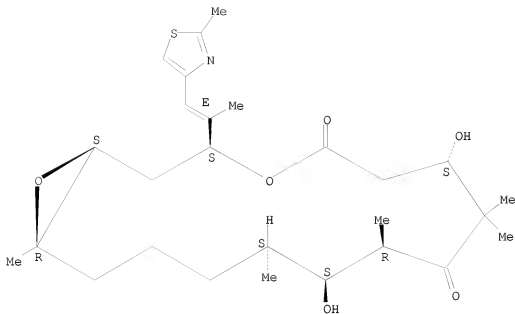
DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N:Y

L2 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2008 ACS on STN

IN 4,17-Dioxabicyclo[14.1.0]heptadecane-5,9-dione, 7,11-dihydroxy-  
8,8,10,12,16-pentamethyl-3-[(1E)-1-methyl-2-(2-methyl-4-thiazolyl)ethenyl]-  
, (1S,3S,7S,10R,11S,12S,16R)-  
MF C27 H41 N O6 S

Absolute stereochemistry. Rotation (-).

Double bond geometry as shown.



\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

=>

---Logging off of STN---

=>

Executing the logoff script...

=> LOG Y

COST IN U.S. DOLLARS

SINCE FILE

TOTAL

FULL ESTIMATED COST

ENTRY

SESSION

13.64

13.85

STN INTERNATIONAL LOGOFF AT 09:03:08 ON 07 JUL 2008

Connecting via Winsock to STN

Welcome to STN International! Enter x:x

LOGINID:SSSPTA1642BJF

PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

\*\*\*\*\* Welcome to STN International \*\*\*\*\*

NEWS 1 Web Page for STN Seminar Schedule - N. America  
NEWS 2 JAN 02 STN pricing information for 2008 now available  
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prophetic substances  
NEWS 4 JAN 28 USPATFULL, USPAT2, and USPATOLD enhanced with new  
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NEWS 6 JAN 28 USGENE now provides USPTO sequence data within 3 days  
of publication  
NEWS 7 JAN 28 TOXCENTER enhanced with reloaded MEDLINE segment  
NEWS 8 JAN 28 MEDLINE and LMEDLINE reloaded with enhancements  
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NEWS 11 FEB 25 IFIREF reloaded with enhancements  
NEWS 12 FEB 25 IMSPRODUCT reloaded with enhancements  
NEWS 13 FEB 29 WPINDEX/WPIDS/WPIX enhanced with ECLA and current  
U.S. National Patent Classification  
NEWS 14 MAR 31 IFICDB, IFIPAT, and IFIUDB enhanced with new custom  
IPC display formats  
NEWS 15 MAR 31 CAS REGISTRY enhanced with additional experimental  
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NEWS 16 MAR 31 CA/Caplus and CASREACT patent number format for U.S.  
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NEWS 17 MAR 31 LPCI now available as a replacement to LDPCI  
NEWS 18 MAR 31 EMBASE, EMBAL, and LEMBASE reloaded with enhancements  
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NEWS 20 APR 15 WPIDS, WPINDEX, and WPIX enhanced with new  
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NEWS 21 APR 28 EMBASE Controlled Term thesaurus enhanced  
NEWS 22 APR 28 IMSRESEARCH reloaded with enhancements  
NEWS 23 MAY 30 INPAFAMDB now available on STN for patent family  
searching  
NEWS 24 MAY 30 DGENE, PCTGEN, and USGENE enhanced with new homology  
sequence search option  
NEWS 25 JUN 06 EPFULL enhanced with 260,000 English abstracts  
NEWS 26 JUN 06 KOREAPAT updated with 41,000 documents  
NEWS 27 JUN 13 USPATFULL and USPAT2 updated with 11-character  
patent numbers for U.S. applications  
NEWS 28 JUN 19 CAS REGISTRY includes selected substances from  
web-based collections  
NEWS 29 JUN 25 CA/Caplus and USPAT databases updated with IPC  
reclassification data  
NEWS 30 JUN 30 AEROSPACE enhanced with more than 1 million U.S.  
patent records  
NEWS 31 JUN 30 EMBASE, EMBAL, and LEMBASE updated with additional  
options to display authors and affiliated  
organizations  
NEWS 32 JUN 30 STN on the Web enhanced with new STN AnaVist  
Assistant and BLAST plug-in  
NEWS 33 JUN 30 STN AnaVist enhanced with database content from EPFULL  
  
NEWS EXPRESS JUNE 27 08 CURRENT WINDOWS VERSION IS V8.3,  
AND CURRENT DISCOVER FILE IS DATED 23 JUNE 2008.  
  
NEWS HOURS STN Operating Hours Plus Help Desk Availability  
NEWS LOGIN Welcome Banner and News Items  
NEWS IPC8 For general information regarding STN implementation of IPC 8

Enter NEWS followed by the item number or name to see news on that

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\* \* \* \* \* STN Columbus \* \* \* \* \*

FILE 'HOME' ENTERED AT 10:09:50 ON 07 JUL 2008

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=> file pctfull
COST IN U.S. DOLLARS          SINCE FILE      TOTAL
                               ENTRY      SESSION
FULL ESTIMATED COST          0.21      0.21
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FILE 'PCTFULL' ENTERED AT 10:10:27 ON 07 JUL 2008  
COPYRIGHT (C) 2008 Univentio

FILE LAST UPDATED: 4 JUL 2008 <20080704/UP>  
FILE COVERS 1978 TO DATE

>>> IMAGES ARE AVAILABLE ONLINE AND FOR EMAIL-PRINTS <<<

>>> NEW FIELD UPTX, FIELD /EW NO LONGER AVAILABLE - SEE HELP CHANGE <<<

```
=> s epothilon?
L1      2484 EPOTHILON?
```

```
=> s l1/ab or l1/ti
        144 EPOTHILON?/AB
        129 EPOTHILON?/TI
L2      159 (EPOTHILON?/AB) OR (EPOTHILON?/TI)
```

```
=> s l2 not py>2001
        817323 PY>2001
L3      53 L2 NOT PY>2001
```

```
=> s combination and l3
        567168 COMBINATION
        264042 COMBINATIONS
        617900 COMBINATION
        (COMBINATION OR COMBINATIONS)
L4      33 COMBINATION AND L3
```

=> d ibib 1-5

```
L4      ANSWER 1 OF 33      PCTFULL  COPYRIGHT 2008 Univentio on STN
ACCESSION NUMBER:          2001092255 PCTFULL  ED 20020826
TITLE (ENGLISH):           EPOTHILONE DERIVATIVES AND METHODS FOR MAKING
                           AND USING THE SAME
TITLE (FRENCH):            DERIVES D'EPOTHILONE, PROCEDES DE PRODUCTION
                           ET METHODES D'UTILISATION
INVENTOR(S):               SANTI, Daniel;
                           FARDIS, Maria;
                           ASHLEY, Gary
PATENT ASSIGNEE(S):        KOSAN BIOSCIENCES, INC.;
                           SANTI, Daniel;
                           FARDIS, Maria;
                           ASHLEY, Gary
DOCUMENT TYPE:             Patent
```

## PATENT INFORMATION:

	NUMBER	KIND	DATE
DESIGNATED STATES	WO 2001092255	A2	20011206
W:	AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW GH GM KE LS MW MZ SD SL SZ TZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE TR BF BJ CF CG CI CM GA GN GW ML MR NE SN TD TG		
PRIORITY INFO.:	US 2000-60/207,655		20000526
	US 2000-60/218,260		20000714
	US 2000-60/231,552		20000911
APPLICATION INFO.:	WO 2001-US15763	A	20010515

L4 ANSWER 2 OF 33 PCTFULL COPYRIGHT 2008 Univention on STN

ACCESSION NUMBER: 2001083800 PCTFULL ED 20020826

TITLE (ENGLISH): PRODUCTION OF POLYKETIDES

TITLE (FRENCH): PRODUCTION DE POLYKETIDES

INVENTOR(S): ARSLANIAN, Robert, L.;

ASHLEY, Gary;

FRYKMAN, Scott;

JULIEN, Bryan;

KATZ, Leonard;

KHOSLA, Chaitan;

LAU, Janice;

LICARDI, Peter, J.;

REGENTIN, Rika;

SANTI, Daniel;

TANG, Li

PATENT ASSIGNEE(S): KOSAN BIOSCIENCES, INC.;

ARSLANIAN, Robert, L.;

ASHLEY, Gary;

FRYKMAN, Scott;

JULIEN, Bryan;

KATZ, Leonard;

KHOSLA, Chaitan;

LAU, Janice;

LICARDI, Peter, J.;

REGENTIN, Rika;

SANTI, Daniel;

TANG, Li

DOCUMENT TYPE: Patent

## PATENT INFORMATION:

	NUMBER	KIND	DATE
DESIGNATED STATES	WO 2001083800	A2	20011108
W:	AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW GH GM KE LS MW MZ SD SL SZ TZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE TR BF BJ CF CG CI CM GA GN GW ML MR NE SN TD TG		
PRIORITY INFO.:	US 2000-09/560,367		20000428
	US 2000-60/232,696		20000914
	US 2000-60/257,517		20001221

US 2001-09/825,856 20010403  
 US 2001-09/825,876 20010403  
 US 2001-60/269,020 20010413  
 APPLICATION INFO.: WO 2001-US13793 A 20010426

L4 ANSWER 3 OF 33 PCTFULL COPYRIGHT 2008 Univentio on STN  
 ACCESSION NUMBER: 2001081341 PCTFULL ED 20020826  
 TITLE (ENGLISH): 9-OXA-EPOTHILON DERIVATIVES, METHOD FOR THE  
 PRODUCTION AND USE THEREOF IN PHARMACEUTICAL  
 PREPARATIONS  
 TITLE (FRENCH): DERIVES DE 9-OXA-EPOTHILONE, LEUR PROCEDE DE  
 PRODUCTION ET LEUR UTILISATION PHARMACEUTIQUE  
 INVENTOR(S): SCHWEDE, Wolfgang;  
 KLAR, Ulrich;  
 SKUBALLA, Werner;  
 BUCHMANN, Bernd;  
 HOFFMANN, Jens;  
 LICHTNER, Rosemarie  
 PATENT ASSIGNEE(S): SCHERING AKTIENGESSELLSCHAFT;  
 SCHWEDE, Wolfgang;  
 KLAR, Ulrich;  
 SKUBALLA, Werner;  
 BUCHMANN, Bernd;  
 HOFFMANN, Jens;  
 LICHTNER, Rosemarie  
 DOCUMENT TYPE: Patent  
 PATENT INFORMATION:

NUMBER	KIND	DATE
WO 2001081341	A2	20011101

DESIGNATED STATES  
 W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU  
 CZ DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS  
 JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN  
 MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR  
 TT TZ UA UG US UZ VN YU ZA ZW GH GM KE LS MW MZ SD SL  
 SZ TZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH CY DE  
 DK ES FI FR GB GR IE IT LU MC NL PT SE TR BF BJ CF CG  
 CI CM GA GN GW ML MR NE SN TD TG  
 PRIORITY INFO.: DE 2000-100 20 899.1 20000420  
 APPLICATION INFO.: WO 2001-EP4551 A 20010419

L4 ANSWER 4 OF 33 PCTFULL COPYRIGHT 2008 Univentio on STN  
 ACCESSION NUMBER: 2001073103 PCTFULL ED 20020822  
 TITLE (ENGLISH): PREPARATION OF EPOTHILONE INTERMEDIATES  
 TITLE (FRENCH): PREPARATION D'INTERMEDIAIRES D'EPOTHILONE  
 INVENTOR(S): VITE, Gregory, D.;  
 KIM, Soong-Hoon;  
 HOEFLE, Gerhard  
 PATENT ASSIGNEE(S): BRISTOL-MYERS SQUIBB COMPANY;  
 VITE, Gregory, D.;  
 KIM, Soong-Hoon;  
 HOEFLE, Gerhard  
 DOCUMENT TYPE: Patent  
 PATENT INFORMATION:

NUMBER	KIND	DATE
WO 2001073103	A2	20011004

DESIGNATED STATES  
 W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR  
 CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL  
 IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG



MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ  
 TM TR TT TZ UA UG US UZ VN YU ZA ZW GH GM KE LS MW MZ  
 SD SL SZ TZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH  
 CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE TR BF BJ  
 CF CG CI CM GA GN GW ML MR NE SN TD TG

PRIORITY INFO.: US 2000-60/191,975 20000324  
 APPLICATION INFO.: WO 2001-US9620 A 20010323

L4 ANSWER 5 OF 33 PCTFULL COPYRIGHT 2008 Univentio on STN  
 ACCESSION NUMBER: 2001070716 PCTFULL ED 20020822  
 TITLE (ENGLISH): A PROCESS FOR THE PREPARATION OF EPOTHILONE  
 ANALOGS AND INTERMEDIATES  
 TITLE (FRENCH): PREPARATION D'ANALOGUES ET D'INTERMEDIAIRES D'  
 EPOTHILONE

INVENTOR(S): LI, Wen, Sen;  
 THORNTON, John, E.;  
 GUO, Zhenrong;  
 SWAMINATHAN, Shankar;  
 PATENT ASSIGNEE(S): MCCONLOGUE, Gary, W.  
 BRISTOL-MYERS SQUIBB COMPANY;  
 LI, Wen, Sen;  
 THORNTON, John, E.;  
 GUO, Zhenrong;  
 SWAMINATHAN, Shankar;  
 MCCONLOGUE, Gary, W.

DOCUMENT TYPE: Patent  
 PATENT INFORMATION:

NUMBER	KIND	DATE
WO 2001070716	A1	20010927

DESIGNATED STATES  
 W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR  
 CU CZ DE DK DM EE ES FI GB GD GE GH GM HR HU ID IL IN  
 IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK  
 MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM  
 TR TT TZ UA UG US UZ VN YU ZA ZW GH GM KE LS MW MZ SD  
 SL SZ TZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH CY  
 DE DK ES FI FR GB GR IE IT LU MC NL PT SE TR BF BJ CF  
 CG CI CM GA GN GW ML MR NE SN TD TG

PRIORITY INFO.: US 2000-09/528,526 20000320  
 APPLICATION INFO.: WO 2001-US7749 A 20010312

=> d his

(FILE 'HOME' ENTERED AT 10:09:50 ON 07 JUL 2008)

FILE 'PCTFULL' ENTERED AT 10:10:27 ON 07 JUL 2008

L1 2484 S EPOTHILON?  
 L2 159 S L1/AB OR L1/TI  
 L3 53 S L2 NOT PY>2001  
 L4 33 S COMBINATION AND L3

=> s l4 and (taxol or paclitaxel)  
 9622 TAXOL  
 272 TAXOLS  
 9705 TAXOL  
 (TAXOL OR TAXOLS)  
 10390 PACLITAXEL  
 72 PACLITAXELS  
 10392 PACLITAXEL  
 (PACLITAXEL OR PACLITAXELS)

L5 29 L4 AND (TAXOL OR PACLITAXEL)

=> s 15 and Her?  
988529 HER?

L6 29 L5 AND HER?

=> s 15 and (HER2 or HER-2)  
4722 HER2  
118696 HER  
1043 HERS  
119313 HER  
(HER OR HERS)

1276185 2  
3260 HER-2  
(HER(W)2)

L7 1 L5 AND (HER2 OR HER-2)

=> d ibib abs

L7 ANSWER 1 OF 1 PCTFULL COPYRIGHT 2008 Univentio on STN  
ACCESSION NUMBER: 1999002514 PCTFULL ED 20020515  
TITLE (ENGLISH): EPOTHILONE DERIVATIVES  
TITLE (FRENCH): DERIVES D'EPOTHILONE  
INVENTOR(S): VITE, Gregory, D.;  
BORZILLERI, Robert, M.;  
KIM, Soong-Hoon;  
JOHNSON, James, A.

PATENT ASSIGNEE(S): BRISTOL-MYERS SQUIBB COMPANY  
LANGUAGE OF PUBL.: English  
DOCUMENT TYPE: Patent  
PATENT INFORMATION:

NUMBER	KIND	DATE
-----		
WO 9902514	A2	19990121

DESIGNATED STATES

W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE  
ES FI GB GE GH GM GW HU ID IL IS JP KE KG KP KR KZ LC  
LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU  
SD SE SG SI SK SL TJ TM TR TT UA UG UZ VN YU ZW GH GM  
KE LS MW SD SZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE  
CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE BF BJ  
CF CG CI CM GA GN ML MR NE SN TD TG

PRIORITY INFO.: US 1997-60/051,951 19970708  
US 1997-60/067,524 19971204  
APPLICATION INFO.: WO 1998-US12550 A 19980616

ABEN The present invention relates to compounds of formula (I), Q is selected from the group consisting of (II), G is selected from the group consisting of alkyl, substituted alkyl, substituted or or unsubstituted aryl, heterocyclo, (III), W is O or NR15; X is O or H,H; Y is selected from the group consisting of O; H,OR16; OR17,OR17; NOR18; H,NOR19; H,NR20R21; H,H; or CHR22; OR17OR17 can be a cyclic ketal; Z1 and Z2 are selected from the group consisting of CH2, O, NR23, S or SO2, wherein only one of Z and Z2 is a heteroatom; B1 and B2 are selected from the group consisting of OR24, or OCOR25, or 2CNR26R27; when B1 is H and Y is OH, H they can form a six-membered ring ketal or acetal; D is selected from the group consisting of NR28R29, NR30COR31 or saturated heterocycle R1, R2, R3, R4, R5, R6, R7, R13, R14, R18, R19, R20, R21, R22, R26 and R27 are

selected from the group H, alkyl, substituted alkyl, or aryl and when R1 and R2 are alkyl can be joined to form a cycloalkyl; R3 and R4 are alkyl can be joined to form a cycloalkyl; R9, R10, R16, R17, R24, R25, and R31 are selected from the group H, alkyl, or substituted alkyl; R8, R11, R12, R28, R30, R32, R33, and R30 are selected from the group consisting of H, alkyl, substituted alkyl, aryl, substituted aryl, cycloalkyl, or heterocyclo; R15, R23 and R29 are selected from the group consisting of H, alkyl, substituted alkyl, aryl, substituted aryl, cycloalkyl, heterocyclo, R32C=O, R33SO2, hydroxy, O-alkyl or O-substituted alkyl, the pharmaceutically acceptable salts thereof and any hydrates, solvates or geometric, optical and stereoisomers thereof, with the proviso that compounds wherein: W and X are both O; and R1, R2, R7 are H; and R3, R4, R6, are methyl; and R8, is H or methyl; and Z1, and Z2, are CH2; and G is 1-methyl-2-(substituted-4-thiazolyl)ethenyl; and Q is as defined above are excluded.

ABFR La presente invention concerne des composés de la formule (I) dans laquelle Q est sélectionné dans le groupe constitué par le groupement (II); G est sélectionné dans le groupe constitué par alkyle, alkyle substitué, aryle substitué ou insubstitué, heterocyclo, le groupement (III); W est O ou NR15; X est O ou H; Y est sélectionné dans le groupe constitué par O; H, OR16; OR17, OR17; NOR18; H, NOR19; H, NR20R21; H; ou CHR22; OR17, OR17 pouvant être un cétal cyclique; Z1 et Z2 sont sélectionnés dans le groupe constitué par CH2, O, NR23, S ou SO2, dans lequel seuls Z et Z2 sont un hétéroatome; B1 et B2 sont sélectionnés dans le groupe constitué par OR24 ou OCOR25 ou OCNR26R27; et peuvent former ensemble un noyau cétal ou acétal à six chaînons si B1 est H et Y est OH, H; D est sélectionné dans le groupe constitué par NR28R29, NR30COR31 ou un hétérocycle saturé. R1, R2, R3, R4, R5, R6, R7, R13, R14, R18, R19, R20, R21, R22, R26 et R27 sont sélectionnés dans le groupe constitué par H, alkyle, alkyle substitué ou aryle, et peuvent former ensemble un cycloalkyle si R1 et R2 ou R3 et R4 sont alkyle; R9, R10, R16, R17, R24, R25 et R31 sont sélectionnés dans le groupe constitué par H, alkyle ou alkyle substitué; R8, R11, R12, R28, R30, R32, R33 et R30 sont sélectionnés dans le groupe constitué par H, alkyle, alkyle substitué, aryle, aryle substitué, cycloalkyle ou heterocyclo; R15, R23 et R29 sont sélectionnés dans le groupe constitué par H, alkyle, alkyle substitué, aryle, aryle substitué, cycloalkyle ou heterocyclo, R32C=O, R33SO2, hydroxy, O-alkyle ou O-alkyle substitué, leurs sels pharmaceutiquement acceptables ou leurs éventuels hydrates, solvates ou isomères géométriques, optiques, ou stéréoisomères, à condition que soient exclus les composés dans lesquels W et X sont tous deux O; et R1, R2 et R7 sont H; et R3, R4 et R6 sont méthyle; et R8 est H ou méthyle; et Z1 et Z2 sont CH2; et G est

1-methyl-2-(substitue-4-thiazolyl)ethenyle; et Q est tel que defini  
ci-dessus.

=> d his

(FILE 'HOME' ENTERED AT 10:09:50 ON 07 JUL 2008)

FILE 'PCTFULL' ENTERED AT 10:10:27 ON 07 JUL 2008

L1 2484 S EPOTHILON?  
L2 159 S L1/AB OR L1/TI  
L3 53 S L2 NOT PY>2001  
L4 33 S COMBINATION AND L3  
L5 29 S L4 AND (TAXOL OR PACLITAXEL)  
L6 29 S L5 AND HER?  
L7 1 S L5 AND (HER2 OR HER-2)

=> s l6 and (HER2 or HER-2)

4722 HER2  
118696 HER  
1043 HERS  
119313 HER  
(HER OR HERS)  
1276185 2  
3260 HER-2

(HER(W)2)  
L8 1 L6 AND (HER2 OR HER-2)

=> s l5 and (HER2 or HER-2)

4722 HER2  
118696 HER  
1043 HERS  
119313 HER  
(HER OR HERS)  
1276185 2  
3260 HER-2

(HER(W)2)  
L9 1 L5 AND (HER2 OR HER-2)

=> d ibib abs kwic

L9 ANSWER 1 OF 1 PCTFULL COPYRIGHT 2008 Univentio on STN  
ACCESSION NUMBER: 1999002514 PCTFULL ED 20020515  
TITLE (ENGLISH): EPOTHILONE DERIVATIVES  
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INVENTOR(S): VITE, Gregory, D.;  
BORZILLERI, Robert, M.;  
KIM, Soong-Hoon;  
JOHNSON, James, A.  
PATENT ASSIGNEE(S): BRISTOL-MYERS SQUIBB COMPANY  
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ABEN The present invention relates to compounds of formula (I), Q is selected from the group consisting of (II), G is selected from the group consisting of alkyl, substituted alkyl, substituted or or unsubstituted aryl, heterocyclo, (III), W is O or NR15; X is O or H,H; Y is selected from the group consisting of O; H,OR16; OR17,OR17; NOR18; H,NOR19; H,NR20R21; H,H; or CHR22; OR17OR17 can be a cyclic ketal; Z1 and Z2 are selected from the group consisting of CH2, O, NR23, S or SO2, wherein only one of Z and Z2 is a heteroatom; B1 and B2 are selected from the group consisting of OR24, or OCOR25, or 2CNR26R27; when B1 is H and Y is OH, H they can form a six-membered ring ketal or acetal; D is selected from the group consisting of NR28R29, NR30COR31 or saturated heterocycle R1, R2, R3, R4, R5, R6, R7, R13, R14, R18, R19, R20, R21, R22, R26 and R27 are selected from the group H, alkyl, substituted alkyl, or aryl and when R1 and R2 are alkyl can be joined to form a cycloalkyl; R3 and R4 are alkyl can be joined to form a cycloalkyl; R9, R10, R16, R17, R24, R25, and R31 are selected from the group H, alkyl, or substituted alkyl; R8, R11, R12, R28, R30, R32, R33, and R30 are selected from the group consisting of H, alkyl, substituted alkyl, aryl, substituted aryl, cycloalkyl, or heterocyclo; R15, R23 and R29 are selected from the group consisting of H, alkyl, substituted alkyl, aryl, substituted aryl, cycloalkyl, heterocyclo, R32C=O, R33SO2, hydroxy, O-alkyl or O-substituted alkyl, the pharmaceutically acceptable salts thereof and any hydrates, solvates or geometric, optical and stereoisomers thereof, with the proviso that compounds wherein: W and X are both O; and R1, R2, R7 are H; and R3, R4, R6, are methyl; and R8, is H or methyl; and Z1, and Z2, are CH2; and G is 1-methyl-2-(substituted-4-thiazolyl)ethenyl; and Q is as defined above are excluded.

ABFR La presente invention concerne des composés de la formule (I) dans laquelle Q est sélectionnée dans le groupe constitué par le groupement (II); G est sélectionné dans le groupe constitué par alkyle, alkyle substitué, aryle substitué ou insubstitué, heterocyclo, le groupement (III); W est O ou NR15; X est O ou H,H; Y est sélectionné dans le groupe constitué par O; H,OR16; OR17,OR17; NOR18; H,NOR19; H,NR20R21; H,H; ou CHR22; OR17,OR17 pouvant être un cétal cyclique; Z1 et Z2 sont sélectionnés dans le groupe constitué par CH2, O, NR23, S ou SO2, dans lequel seuls Z et Z2 sont un hétéroatome; B1 et B2 sont sélectionnés dans le groupe constitué par OR24 ou OCOR25 ou 2CNR26R27; et peuvent former ensemble un noyau cétal ou acétal à six chaînons si B1 est H et Y est OH,H; D est sélectionné dans le groupe constitué par NR28R29, NR30COR31 ou un heterocycle saturé. R1, R2, R3,

R4, R5, R6, R7, R13, R14, R18, R19, R20, R21, R22, R26 et R27 sont selectionnees dans le groupe constitue par H, alkyle, alkyle substitue ou aryle, et peuvent former ensemble un cycloalkyle si R1 et R2 ou R3 et R4 sont alkyle; R9, R10, R16, R17, R24, R25 et R31 sont selectionnees dans le groupe constitue par H, alkyle ou alkyle substitue; R8, R11, R12, R28, R30, R32, R33 et R30 sont selectionnees dans le groupe constitue par H, alkyle, alkyle substitue, aryle, aryle substitue, cycloalkyle ou heterocyclo; R15, R23 et R29 sont selectionnees dans le groupe constitue par H, alkyle, alkyle substitue, aryle, aryle substitue, cycloalkyle ou heterocyclo, R32C=O, R33SO2, hydroxy, O-alkyle ou O-alkyle substitue, leurs sels pharmaceutiquement acceptables ou leurs eventuels hydrates, solvates ou isomeres geometriques, optiques, ou stereoisomeres, a condition que soient exclus les composés dans lesquels W et X sont tous deux O; et R1, R2 et R7 sont H; et R3, R4 et R6 sont methyle; et R8 est H ou methyle; et Z1 et Z2 sont CH2; et G est 1-methyl-2-(substitue-4-thiazolyl)ethenyle; et Q est tel que defini ci-dessus.

TIEN EPOTHILONE DERIVATIVES

TIFR DERIVES D'EPOTHILONE

DETD

R  
S Me  
jOH  
N3 ], ]' ] '  
O Me  
O OH O  
I EpothiloneA R=H  
II EpothiloneB R=Me  
have been found to exert microtubule-stabilizing effects similar to  
TAXOL and hence cytotoxic activity against rapidly  
proliferating cells,  
such as, tumor cells or other hyperproliferative cellular disease, see  
.Angew. Chem. Int. Ed. Engl., . . .

The compounds of this invention. are also useful in combination with known anti-cancer and cytotoxic agents and treatments, including radiation. If formulated as a fixed dose, such combination products employ the compounds of this invention within the dosage range described below and the other pharmaceutically active agent within its approved dosage range. Compounds of formula V can be used sequentially with known anticancer or cytotoxic agents and treatment, including radiation when a combination formulation is inappropriate.

Especially useful are cytotoxic drug combinations wherein the second drug chosen acts in a different phase of the cell cycle, e.g. S phase, than the present compounds of. . .

. . .  
Synthase Inhibitors,  
DNA Cross Linking Agents  
Topoisomerase I and II Inhibitors  
DNA Alkylating Agents

Ribonucleoside Reductase Inhibitors  
Cytotoxic Factors e.g. TNF-alpha or  
Growth factor inhibitors e.g. HER 2 receptor MAB's  
The present compounds may exist as multiple optical, geometric,  
and stereoisomers. Included within the present invention are all such  
isomers and. . .

. . .  
potency is  
accomplished following a modified procedure of Swindell, et al., (see  
Swindell, C.S., Krauss, N.E., Horwitz, S.B., and Ringel, I. Biologically  
active taxol analogues with deleted A-ring side chain  
substituents and  
variable C-2' configurations. J. Med. Chem. 34: 1176-1184, 1991). These  
modifications, in part, result. . .

. . .  
cells were incubated at 37' for 72 hours at which time the  
tetrazolium dye, MTS at 333 gg/ml (final concentration), in  
combination  
with the electron coupling agent phenazine methosulfate at 25 gm (final  
concentration) was added. A dehydrogenase enzyme in live cells  
reduces the MTS. . .

=>

---Logging off of STN---

=>

Executing the logoff script...

=> LOG Y

COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	23.30	23.51

STN INTERNATIONAL LOGOFF AT 10:14:45 ON 07 JUL 2008